WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6: C07K 14/72, C12N 15/10, A61K 38/17 (11) International Publication Number:

WO 99/19354

(43) International Publication Date:

22 April 1999 (22.04.99)

(21) International Application Number:

PCT/SE98/01548

(22) International Filing Date:

31 August 1998 (31.08.98)

(30) Priority Data:

9703745-1 9801148-9 14 October 1997 (14.10.97) SE

31 March 1998 (31.03.98) SE

(71) Applicant (for all designated States except US): PHARMACIA & UPJOHN AB [SE/SE]; S-112 87 Stockholm (SE).

(72) Inventors; and

(75) Inventors/Applicants (for US only): BERKENSTAM, Anders [SE/SE]; Långholmsgatan 11, S-117 33 Stockholm (SE). DAHLBERG, Mats [SE/SE]; Sankt Göransgatan 64, S-112 87 Stockholm (SE).

(74) Agents: TANNERFELDT, Agneta et al.; Pharmacia & Upjohn AB, S-112 87 Stockholm (SE).

(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, HR, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

With international search report.

(54) Title: NOVEL VITAMIN D RECEPTOR RELATED POLYPEPTIDES, NUCLEIC ACID SEQUENCE ENCODING THE SAME AND USES THEREOF

(57) Abstract

The present invention relates to novel vitamin D receptor related (VDRR) polypeptides, and formulations containing the same. Nucleic acid sequences encoding the VDRR polypeptides, expression vectors containing such sequences and host cells transformed with such expression vectors are also disclosed, as are methods for the expression of the novel VDRR polypeptides of the invention. The invention further relates to VDRR polypeptides for use as medicaments, and use of substances affecting VDRR signal transduction for the manufacture of medicaments for treating metabolic, proliferative or inflammatory conditions. The present invention also relates to methods for identifying clones encoding a VDRR polypeptide, methods for identifying ligands to a VDRR and methods for identifying substances for treatment of conditions affected by a VDRR polypeptide. More specifically, the novel VDRR polypeptide can be the polypeptide designated VDRRγ, which may be regulated by any small chemical molecule similar in structure to known ligands for nuclear receptors.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	ES	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	Prance	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	I.E	treland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Nige r	VN	Vict Nam
CG	Congo	KE	Kenya	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	zw	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
CZ	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	ш	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		

WO 99/19354 PCT/SE98/01548

NOVEL VITAMIN D RECEPTOR RELATED POLYPEPTIDES, NUCLEIC ACID SEQUENCE ENCODING THE SAME AND USES THEREOF

5 FIELD OF THE INVENTION

10

15

20

25

30

The present invention relates to novel vitamin D receptor related (VDRR) polypeptides. Nucleic acid sequences encoding the same, expression vectors containing such sequences and host cells transformed with such expression vectors are also disclosed, as are methods for the expression of the novel VDRR polypeptides of the invention, and uses thereof.

BACKGROUND OF THE INVENTION

Nuclear hormone receptors is a large group of conditionally regulated transcription factors. These receptors are activated and regulate target gene expression in response to binding a variety of small chemical molecules (ligands) including steroids, vitamin D3, retinoids, eicosanoides (prostanoids), thyroid hormone and cholesterol derivatives.

A growing number of structurally related receptors have been identified for which no ligands yet have been identified. This group of receptors is referred to as orphan nuclear receptors (ONRs). A review of the ONRs can be found in Enmark et al, Mol. Endo., vol. 10, No. 11 (1996) pp. 1293-1307, which is hereby incorporated by reference. The pivotal importance of a number of ONRs for processes such as metabolic homeostasis, cell differentiation and development have been demonstrated both by biochemical and genetic techniques. In addition, several ONRs have also been implicated as key factors in a variety of common diseases and disorders such as diabetes, obesity, inflammatory conditions and proliferative diseases.

Based on these findings it is generally believed that novel ONRs are going to become potential drug targets for therapeutic invention of common diseases. Thus, it is of great importance to identify such receptors.

SUMMARY OF THE INVENTION

The present invention relates to novel vitamin D receptor related (VDRR) polypeptides, and formulations containing the same. Nucleic acid sequences encoding the VDRR polypeptides, expression vectors containing such sequences and host cells transformed with such expression vectors are also disclosed, as are methods for the expression of the novel VDRR polypeptides of the invention. The invention further relates to VDRR polypeptides for use as medicaments, and use of substances affecting VDRR signal transduction for the manufacture of medicaments for treating metabolic, proliferative or inflammatory conditions. The present invention also relates to methods for identifying clones encoding a VDRR 10 polypeptide, methods for identifying ligands to a VDRR and methods for identifying substances for treatment of conditions affected by a VDRR polypeptide. More specifically, the novel VDRR polypeptide can be the polypeptide designated VDRRy, which may be regulated by any small chemical molecule similar in structure to known ligands for nuclear receptors.

BRIEF DESCRIPTION OF THE DRAWINGS

15

- Figure 1 The cDNA sequence encoding the novel nuclear receptor polypeptide 20 vitamin D receptor related gamma (VDRRg) is shown.
 - Figure 2 Evolutionary neighbor-joining tree for VDRRg as given by DBD-HMM alignment.
 - Figure 3 Evolutionary neighbor-joining tree for VDRRg as given by LBD-HMM alignment.
- 25 Figure 4 - The deduced amino acid sequence of VDRRg is shown.
 - Figure 5 Expression of VDRRg in adult human tissues. The numbers on the right hand side, refer to kilobasepairs of the mRNA.
- Figure 6 Vitamin D3 transactivate a GAL4-DBD/VDR-LBD fusion protein but not a GAL4-DBD/VDRRy-LBD fusion protein in transient transfections of CV-1 cells. The 30 number on the left hand side refer to relative luciferase activity of the GAL4-luciferase reporter gene.

WO 99/19354 PCT/SE98/01548

Figure 7 - The cDNA sequence encoding VDRRg-2 with an alternatively spliced 5'-end compared to VDRRg is shown.

- Figure 8 The deduced amino acid sequence of VDRRg-2 is shown.
- Figure 9 Heterodimerization of VDRRg with a retinoid X receptor (RXR) is shown.
- Figure 10 The effect of pregnenolone derivatives as activators of VDRRg are shown.
 - Figure 11 The effect of pregnenolone 16α-carbonitrile (PCN), dexamethasone and an antiprogestin (RU486) as activators of VDRRg are shown.
 - Figure 12 Percent similarity between the new genes VDRRg-1 and VDRRg-2 and the known genes XOR-6. HVDR, CAR-1 and CAR-2.
 - Figure 13 Percent identity between the new genes VDRRg-1 and VDRRg-2 and the known genes XOR-6. HVDR, CAR-1 and CAR-2.

15

20

25

30

10

DETAILED DESCRIPTION OF THE INVENTION

The objects above are met by the present invention, which relates to a mammalian, preferably human, isolated or recombinant nucleic acid comprising a contiguous nucleic acid sequence encoding a vitamin D receptor related (VDRR) polypeptide. The VDRR polypeptide is suitably origin.

In preferred embodiments of the present invention, the nucleic acid encoding the VDRR polypeptide contains a DNA-binding domain (DBD) comprising about 77 amino acids with 9 cysteine residues. The DBD is further characterized by the following amino acid sequence similarity relative to the DBDs of human Vitamin D Receptor (hVDR) and Orphan Nuclear Receptor 1 isolated from *Xenopus laevis* (xONR1 = XOR-6), respectively: (i) at least about 60% amino acid sequence similarity with the DBD of hVDR; and (ii) at least about 65% amino acid sequence similarity with the DBD of xONR1. More particularly, the amino acid sequence similarity relative to the DBDs of hVDR and xONR1, respectively is

(i) about 65% amino acid sequence similarity with the DBD of hVDR; and

WO 99/19354 4 PCT/SE98/01548

(ii) about 71% amino acid sequence similarity with the DBD of xONR1.

In preferred embodiments of the present invention, the nucleic acid encoding the VDRR polypeptide contains a ligand-binding domain (LBD) characterized by the following amino acid sequence similarity, relative to the LBDs of hVDR and xONR1, respectively:

- (i) at least about 30% amino acid sequence similarity with the LBD of hVDR, suitably at least 35% amino acid sequence similarity with the LBD of hVDR; and
 - (ii) at least about 40% amino acid sequence similarity with the LBD of xONR1, suitably at least 45% amino acid sequence similarity with the LBD of xONR1.

More particularly, the amino acid sequence similarity relative to the LBDs of hVDR and xONR1, respectively is

- (i) about 42% amino acid sequence similarity with the LBD of hVDR; and
- (ii) about 54% amino acid sequence similarity with the LBD of xONR1.

20

25

30

- "amino acid sequence similarity" refers to: 100x Consensus Lenght divided by Consencus Length + Mismatsches + Gaps.
- The term amino acid sequence identity can also be used. Amino acid sequence identity is calculated by comparing the absolute amino acid residue identity. In Figure 13 the amino acid sequence identity between the new genes VDRRg-1 and VDRRg-2 and the known genes are shown.

In particularly preferred embodiments, the nucleic acid sequences of the present invention are substatially the same as those given in Fig. 1 or Fig. 7, the same or alleles thereof.

The present invention also relates to a nucleic acid probe for the detection of a nucleic acid sequence encoding a VDRR polypeptide in a sample. Suitably, the probe comprises at least 14 contiguous nucleotides, and preferably at least 28 contiguous nucleotides, of the nucleic acid sequences given in Fig. 1 or Fig. 7. The nucleic acid probe can be used in a method for identifying clones encoding a VDRR polypeptide, wherein the method comprises screening a genomic or cDNA library with the probe under low stringency hybridization conditions, and identifying those clones which display a substantial degree of hybridization to said probe.

The present invention further relates to an isolated or recombinant VDRR polypeptide. The polypeptide can be full-length, at which the sequence of amino acids is identical to WO 99/19354 5 PCT/SE98/01548

the corresponding sequence found in mammals in general, and in human beings in particular. In the present invention, the polypeptide can also be a truncated, extended or mutated form of the full-length polypeptide. Truncated and extended forms relate to VDRR polypeptides where one or more amino acids are missing or have been added, respectively, at the N terminal end of the polypeptide chain. Mutated forms relate to VDRR polypeptides where one or more amino acid has been substituted by another amino acid. Suitably, the isolated or recombinant VDRR polypeptide exhibits the amino acid sequences given in Fig. 4 or Fig. 8.

5

10

15

20

25

The N-terminal sequence of the present nucleic acids encoding VDRR polypeptides, as well as the amino acid sequence of the present VDRR polypeptides, may vary. Thus, various N-terminal isoforms are envisaged, e.g. any of $\alpha 1$, $\alpha 2$, $\beta 1$, $\beta 2$, $\beta 3$, $\beta 4$, $\gamma 1$ or $\gamma 2$ as disclosed in Fig. 7B of Transcription Factors 3: nuclear receptors, Protein Profile, vol. 2, issue 11 (1995), pp. 1173-1235. This review of nuclear receptors generally is hereby incorporated by reference. More specifically, Vitamin D receptors and related orphans, e.g. ONR1, are discussed at p. 1191-1992.

The present invention further relates to pharmaceutical formulations comprising an isolated or recombinant VDRR polypeptide, and one or more therapeutically acceptable excipients. Examples of excipients that can be used are carbohydrates, e.g. monosaccharides, disaccharides and sugar alcohols, such as saccharose and sorbitol. Further examples include amino acids, e.g. histidine and arginine, surfactants, e.g. polyoxyethylene sorbitan fatty acid esters, inorganic salts, e.g. sodium chloride and calcium chloride, and complexing agents, e.g. EDTA and citric acid.

The present formulation can be in the form of an aqueous solution ready-for-use, or dried, particularly lyophilized. In the latter case, the formulation is reconstituted with a liquid, e.g. sterile water or saline, before use.

The present invention further relates to an expression vector comprising an isolated or recombinant nucleic acid, the nucleic acid comprising a contiguous nucleic acid sequence encoding a Vitamin D receptor related (VDRR) polypeptide. The invention also relates to a cell containing such an expression vector.

WO 99/19354 6 PCT/SE98/01548

The present invention further relates to a cell containing the claimed nucleic acid, the nucleic acid comprising a contiguous nucleic acid sequence encoding a Vitamin D receptor related (VDRR) polypeptide.

The present invention further relates to a process for recombinant production of a VDRR polypeptide, by expressing the claimed isolated or recombinant contiguous nucleic acid sequence encoding a Vitamin D receptor related (VDRR) polypeptide in a suitable host cell, preferably an eukaryotic cell.

5

10

15

20

25

30

The present invention further relates to method for identifying a ligand to a VDRR, e.g. by a cell-based reporter assay, transgenic-animal reporter assay or *in vitro*-binding assay. It also relates to a method for identifying a substance for treatment of a condition affected by a VDRR polypeptide, comprising screening for an agonist or an antagonist of VDRR polypeptide signal transduction to be used for treating metabolic, proliferative or inflammatory conditions.

The present invention further relates to a VDRR polypeptide for use as a medicament, as well as use of a substance affecting VDRR signal transduction for the manufacture of a medicament for treating metabolic, proliferative or inflammatory conditions.

More particularly, the present invention can be used for the manufacture of medicaments for treating obesity, diabetes, anorexia, lipoprotein defects, hyperlipidemia, hypercholeste-remia or hyperlipoproteinemia. The present invention can be used also for the manufacture of medicaments for treating osteoporosis, rheumatoid artritis, benign and malign tumors, hyperproliferative skin disorders or hyperparathyroidism.

The present invention further relates to a method for treating metabolic, proliferative or inflammatory conditions by introducing into a mammal a nucleic acid vector encoding for expression of a VDRR polypeptide. The nucleic acid vector is capable of transforming a cell *in vivo* and expressing said polypeptide in said transformed cell.

The present invention further relates to a method for treatment of a metabolic, proliferative or inflammatory condition by administration of a therapeutically effective amount of a substance affecting VDRR signal transduction, specifically a VDRR polypeptide.

In the present invention, the term "isolated" in connection with VDRR polypeptides or nucleic acids encoding the same, relates to nucleic acids or polypeptides that have been isolated from a natural source, e.g. the liver, small intestine or colon of a human being. The

WO 99/19354 7 PCT/SE98/01548

isolated VDRR polypeptides or nucleic acids of the present invention are unique in the sense that they are not found in a pure or separated form in nature. Use of the term "isolated" indicates that a naturally occurring sequence has been removed from its normal cellular environment. Thus, the sequence may be in a cell-free environment or in a different cellular environment. The term does not imply that the sequence is the only nucleic acid or amino acid sequence present, but that it is the predominant nucleic acid or amino acid sequence present. Furthermore, the nucleic acid or polypeptide should be essentially free of non-amino acid or non-nucleic acid material naturally associated with the respective product. In this context, essentially free relates to more than 80%, suitably more than 90%, and preferably more than 95% purity.

The term "sustantially the same "when referring to the nucleic acid sequences in Fig 1 or Fig 7 and when referring to the amino acid sequences in Fig. 4 or Fig. 8 means that they are derived from the sequences given in the figures and have the same function as those.

10

15

20

25

30

The inventors of the present invention, have surprisingly isolated a novel nucleic acid sequence, and a polypeptide encoded by said nucleic acid sequence. Thus, a novel cDNA encoding a polypeptide designated VDRRγ has been cloned and characterized. This polypeptide is, based on amino acid sequence similarity, a novel member of the nuclear (hormone) receptor supergene family. Hidden Markov Models (HMMs) in combination with phylogenetic analysis such as neighbor-joining tree methods and other statistical algorithms shows that VDRRγ belong to a sub-family of vitamin D receptors (VDRs) and a VDR-like receptor from *Xenopus laevis* designated xONR1 (see Smith et al., Nucl. Acids Res., 22 (1994), No. 1, pp. 66-71) or XOR-6 as in WO96/22390. The VDRRγ, therefore, is one member of a family of Vitamin D receptor related (VDRR) polypeptides.

The degree of amino acid similarity in the DBD and LBD of VDRRg as compared to the

most closely related receptors XOR-6, hVDR and CAR (see WO 93/17041) is similar to the relationship between other distinct, but related nuclear receptors. (See Fig.12). The thyroid hormone (TRb) and retinoic acid receptor (RARb) are approximately 60% and 40% identical at the amino acid level in the DBD and LBD, respectively. By comparison, the closely related but unique genes encoding human RARa and RARb nuclear receptors are 97% and 82% identical in the DBD and LBD, respectively.

WO 99/19354 8 PCT/SE98/01548

As recognized by those skilled in the art of nuclear receptors, the DBD displays the highest degree of conservation (amino acid identity) both between different nuclear receptors (paralogous) and between identical receptors from different species (orthologues). The two "zink-fingers" in the DBD are generated by two evolutionary conserved amino acid motifs Cys-X2-Cys-X13-Cys-X2-Cys (amino-terminal or first zink-finger) and Cys-Xn-Cys-X9-Cys-X2-Cys (carboxy-terminal or second zink-finger) in which two pairs of cysteins chelate on zink ion. The vast majority of nuclear receptors have five amino acid residues between the firs two Cys residues in the second zink-finger (Cys-X5-Cys-X9-Cys-X2-Cys) see Gronemeyer and Laudet (Protein Profile 1995, 2, issue 11) for details. The today only known exception to this role are the PPARs which have three amino acid (Cys-X3-Cys-X9-Cys-X2-Cys) residues and the TLL group of receptors which have seven (Cys-X7-Cys-X9-Cys-X2-Cys). Thus another feature which is characteristic of the novel VDRRg polypeptide described herein is that the number of amino acid residues in this part of the DBD is six (Cys-X6-Cys-X9-Cys-X2-Cys) as shown in Figs.4 and 8. Today, the only other nuclear receptor like sequences found in the TREMBLE data base with the same number of amino acid residues between the two cys residues are two sequences (Q20097 and Q18155) from the worm C. elegans (Q20097 and Q18155). However, the entire DBD of these putative C. elegans nuclear receptors are only distantly related to the DBD of VDRRg. Taken together, the comparison of the DBD and LBD of the nuclear receptor VDRRg described herein (See Fig.12), clearly demonstrate that this receptor is a novel member of the nuclear receptor super-gene family which is distinct from other known nuclear receptors that are most closely related to the VDRRg including ONR-1 (in Smith et al., 1994, Nucleic Acids Res., 22, pp66-71) or XOR-6 (in WO 96/22390), hVDR and CAR (WO 93/17041). This finding, in combination with the highly restricted expression pattern we observe for human VDRRy (liver, small intestine and mucosa of colon) and in analogy to other nuclear receptors exhibiting a tissue specific expression pattern such as the peroxisome pro-liferatoractivated receptors (PPARs) - suggest that VDRRy performs important physiolo-gical functions in liver, small intestine and colon. Accordingly, VDRRy is likely to be an important sensor of key metabolic pathways affecting lipid, carbohydrate or amino acid

metabolism/homeostasis. In addition, the highly selective tissue specific expression pattern

10

15

20

25

30

WO 99/19354 9 PCT/SE98/01548

suggest that VDRRy may participate in cellular differentiation and development of these tissues.

An additional human VDRRγ cDNA with an alternatively spliced 5'-end has been identified (see Fig. 7). The VDRRγ cDNAs are thus able to encode at least one alternative N-terminal variant (Fig. 8) in addition to the VDRRγ polypeptide shown in Fig. 4. In analogy to other members of the nuclear receptor supergene family such as RORα and RARα these N-terminal isoforms of VDRRγ may specify different functions including DNA-binding specificity and/or promoter specific activation (Gronemeyer and Laudet, 1995).

5

10

15

20

25

In the present specification, the term VDRRγ relates to the various polypeptides corresponding to the differentially spliced VDRRγ cDNAs including VDRRγ-1 and VDRRγ-2. However, when reference is made to Fig. 1 and Fig. 4, VDRRγ cDNA and VDRRγ relates specifically to VDRRγ-1 cDNA and VDRRγ-1, respectively. In the same way, when reference is made to Fig. 7 and Fig. 8, VDRRγ cDNA and VDRRγ relates specifically to VDRRγ-2 cDNA and VDRRγ-2, respectively.

In contrast to the VDRRγ-2 cDNA, the VDRRγ-1 cDNA does not contain a classical AUG initiation codon but instead may initiate at an alternative CUG codon. This putative non-AUG start site is located in a favorable sequence context for efficient initiation from alternative start sites and is in frame with the entire open reading frame and preceded by a stop codon.

Taken together, the VDRRs in general, and more specifically the VDRRγ, may be important in

- 1) metabolic diseases such as obesity, diabetes (type I and II), lipoprotein disorders,
- 2) proliferative conditions such as tumors (benign and malignant) of the small intestine and colon,
- 3) ulcero-inflammatory diseases of small intestine and colon such as Crohn's disease and ulcerative colitis, and
- 4) congenital anomalies of small intestine and colon.

The high amino acid sequence identity of VDRRy with the VDR both in the DNAbinding domain (DBD) and ligand-binding domain (LBD) indicate that these two receptors WO 99/19354 10 PCT/SE98/01548

5

10

15

20

25

30

may also have overlapping yet distinct functional characteristics. In analogy, retinoic acid receptors (RARs) and retinoid X receptors (RXRs) have similar amino acid sequence identities in the DBD and LBD region as the VDR and VDRRγ. RARs and RXRs have been shown to have distinct functional similarities such that both receptors bind 9-cis retinoic acid and have overlapping DNA-binding specificities and accordingly regulate overlapping gene networks. Based on these findings, VDRRγ may be regulated by small chemical molecules similar in structure to known ligands for nuclear receptors but not necessarily identical to ligands for the 1α, 25-dihydroxy vitamin D3 receptor. Furthermore, VDRRγ may regulate vitamin D3 responsive gene networks by binding to a Vitamin D responsive element (VDRE)-like DNA sequence. In the present application, the 1α, 25-dihydroxy vitamin D3 receptor is abbreviated as the Vitamin D receptor (VDR).

In the present invention, the substance affecting VDRR signal transduction can be any small chemical molecule of natural or synthetic origin, e.g. a carbohydrate such as an aromatic compound. The small molecule may have a molecular weight in the range of from about 100 up to about 500 Da. Suitably, the small chemical molecule has a molecular weight in the range of from 200 up to 400 Da. Preferably, the small chemical molecule has a molecular weight of about 300 Da.

The human VDRRγ polypeptides, including VDRRγ-1 and VDRRγ-2, have been shown to be activated e.g. by pregnenolones and estradiol (weakly), but not by certain other steroid hormones such as cortisol, aldosterone, progesterone and estrogen, and most likely not by progestines and glucocorticoids. Thus, human VDRRγ is not activated by pregnenolone 16α-carbonitrile (PCN), a glucocorticoid antagonist. For this reason, human VDRRγ can also be designated human pregnenolone activated (nuclear) receptors (hPAR). Information about pregnenolone can be found e.g. in the Merck Index, 11th ed., Merck & Co., Inc. Rahway, N.J., USA, p. 7735, 1989.

Activators for human VDRRγ polypeptides, including VDRRγ-1 and VDRRγ-2, (hPAR-1 and hPAR-2, respectively), include but are not limited to pregnenolones, such as pregnane-ones, pregnane-diones, pregnane-triones, and pregnane-diols, and androstanes, such as androstane-ols, and androstane-diols. Suitably, the pregnenolones are non-planar, particularly 5β-pregnanes.

Specific examples of activators and possibly ligands for human VDRR γ polypeptides, including VDRR γ -1 and VDRR γ -2, are the following compounds, which are marketed by Sigma-Aldrich of Sweden:

- i) 5β-pregnane-3,20-dione
- 5 ii) 3α-hydroxy-5β-pregnane-11,20-dione methanesulphonate
 - iii) 5β-pregnane-3α,20β-diol
 - iv) pregnenolone
 - v) Pregn-4-eno[16,17- δ][2]isoxazolline-3,20-dione, 6α -methyl-3'-phenyl-, ethyl ether solvate
- vi) Pregna-1,4,9(11)-triene-3,20-dione, 21-[4-[6-methoxy-2-(4-morpholinyl)-4-pyrimidinyl]-1-piperazinyl]-16-methyl-, (16α)
 - vii) Estran-3-ol, 17-[[[3-(trifluoromethyl)phenyl]methyl]amino]-, (E)-2-butenedioate (1:1) (salt)
 - viii) 9α-Fluoro-5α-androstane-11β,17β-diol
- 15 ix) Spiro[5α-androstane-3,2'-benzothiazolin]-11-one, 17β-hydroxy-17-methyl
 - x) Spiro[pregnane-3,2'-thiazolidine]-4'-carboxylic acid, 11α -hydroxy-20-oxo-, sodium salt
 - xi) 17β-Dimethylamino-17-ethynyl-5α-androstane-11β-ol
 - xii) 6β-Hydroxy-3,5-cyclo-5α-pregnan-20-one, nitrite
- 20 xiii) 3α-Hydroxy-5β-pregnane-11,20-dione, acetate, 20-O-(methylsulfonyl)-oxime
 - xiv) 17α -Methyl- 5α -androstane- 11β ,17-diol
 - xv) 5β-Pregnane-3,11,20-trione, trioxime
 - xvi) 3α -Hydroxy- 5β -pregnane-11,20-dione, 20-hydazone with hydrazide of 1-(carboxymethyl) pyridinium chloride.
- A possible use of a VDRRg antagonist, could be a synergistic co-administration of the VDRRg antagonist together with other drugs such as, but not limited to, HIV protease inhibitors and cyclosporin to inhibit the expression of CYP3A4 and thus increase the bioavailability of drugs with poor pharmacokinetics due to CYP3A4 metabolism.

 Genes coding for polypeptides, such as human vitamin D receptor related gamma
- 30 (hVDRRg), may be cloned by incorporating a DNA fragment coding for the polypeptide into a recombinant DNA vehicle, e.g. a vector, and transforming suitable prokaryotic or

WO 99/19354 12 PCT/SE98/01548

eukaryo-tic host cells. Such recombinant DNA techniques are well known and e.g. described in Methods in Enzymology, Academic Press, San Diego, CA, USA (1994), vols. 65 and 68 (1979), and vols. 100 and 101 (1983).

The host cells for use in the present invention can be prokaryotic or eukaryotic, preferably eukaryotic cells. Suitable eukaryotic host cells include but are not limited to cells from yeast, e.g. Saccharomyces, insect cells and mammalian cells such as Chinese Hamster Ovary (CHO), Baby Hamster Kidney (BHK), COS and the like. Suitable prokaryotic host cells include but are not limited to cells from Enterobacteriacea, e.g. E. coli, Bacillus and Streptomyces.

10

15

20

25

30

5

EXAMPLES

The following Examples are provided for purposes of illustration only and are not to be construed as in any way limiting the scope of the present invention, which is defined by the appended claims.

EXAMPLE 1

Identification and isolation of human VDRRg cDNA

Expressed Sequence Tag (EST) databases were screened for nuclear receptor related sequences with a DNA-binding domain (DBD) profile of nuclear receptors. This search profile was created by multiple alignment of a selected set of nuclear receptor sub-domains followed by a statistical calculation to obtain a so called Hidden Markov Model (HMM) of different subfamily members of the nuclear receptor supergene family. The cDNA of one of the nuclear receptor related EST sequences identified (Incyte clone no 2211526) was analyzed in detail by sequencing. After DNA sequencing of the entire Incyte cDNA clone (approximately 2200 basepairs) the clone was found to encode a putative ligand-binding domain (LBD) with 54% and 44% similarity to xONR-1 and to the vitamin D receptor (VDR), respectively. The cDNA of the Incyte clone was not full-length and did not encode a sequence corresponding to a complete DBD.

WO 99/19354 13 PCT/SE98/01548

5'-RACE (rapid amplification of cDNA ends) of random primed cDNA from human liver RNA (InVitrogen) followed by cloning and DNA sequencing showed that the 5'-part of the cDNA corresponding to the Incyte clone encoded a DBD characteristic for nuclear receptors and with 71% and 65% sequence similarity to xONR-1 and VDR, respectively. Multiple alignments in combination with evolutionary neighbor-joining tree analysis placed the polypeptide encoded by the cDNA (specified in Fig. 1) in the group of VDRs (Figs. 2 and 3) and was named human vitamin D receptor related gamma (VDRRg). The deduced amino acid sequence of VDRRg is given in Fig. 4.

10 EXAMPLE 2

5

15

20

25

30

Expression of VDRRg mRNA in human tissues

Multiple tissue northern blots (Clontech) was used to determine the expression pattern of VDRRg in adult human tissues. As shown in Fig. 5, VDRRg is abundantly expressed in small intestine, mucosal lining of colon and liver but not in several other tissues including spleen, thymus, prostate, testis, ovary, peripheral blood leukocytes, heart, brain, placenta, lung, skeletal muscle, kidney and pancreas. To investigate if VDRRγ was expressed at lower levels in any of the other tissues examined, the filter was exposed for an extended time (one week as compared to overnight). Even after this prolonged exposure (data not shown), expression could still only be detected in the same tissues and not in any of the other tissues examined. The restricted expression pattern of VDRRg suggest that this receptor is likely to have an important regulatory function in liver and intestine.

EXAMPLE 3

Transient transfections of GAL4-DBD/VDRRy-LBD fusion protein using Vitamin D3

Transient transfections were performed to analyze if vitamin D3 activate the VDRRy polypeptide. To this end, transient co-transfections of CV-1 cells were performed with expression plasmids encoding fusion proteins of the GAL4-DBD fused to the LBD of either the VDR or the VDRR together with a reporter-plasmid containing five GAL4 responsive elements upstream of the luciferase gene. After transfection, cells were treated with vehicle (DMSO) alone or with vitamin D3 for 48 hours followed by harvesting of the cells and measurement of the luciferase activity in cell extracts. As shown in Fig. 6, vitamin D3 (1

WO 99/19354 14 PCT/SE98/01548

μM) transactivate the GAL4-DBD/VDR-LBD but not the corresponding GAL4-DBD/-VDRRγ-LBD polypeptide under these conditions. This indicates that the two receptors may have distinct ligand-binding specificities.

5 EXAMPLE 4

Identification and isolation of human VDRRy cDNAs encoding multiple N-terminal isoforms

5'-RACE (see Example 1) of cDNA from human liver RNA followed by cloning and DNA sequencing identified an additional human VDRRγ cDNA with alternatively spliced 5'-end (see Fig. 7). The VDRRγ cDNAs are thus able to encode at least one alternative N-terminal variant (Fig. 8) in addition to the VDRRγ polypeptide shown in Fig. 4. The polypeptides disclosed in Fig. 4 and Fig. 8 which correspond to the differentially spliced VDRRγ cDNAs are designated as VDRRγ-1 and VDRRγ-2, respectively.

15

20

25

30

10

EXAMPLE 5

VDRRy heterodimerise with RXR and bind to direct repeats (DRs) spaced by three or four nucleotides

Expression plasmids containing VDRRγ or RXRβ cDNAs were transcribed using T7 polymerase and translated *in vitro* in TNT reticulocyte lysates (Promega, Madison, WI, USA). To investigate the DNA-binding specificity of VDRRγ a native gel mobility assay was employed essentially as described (Berkenstam et al., Cell, 69, 401-412, 1992) in which *in vitro* translated VDRRγ was incubated in the presence or absence of *in vitro* translated RXRβ with different 32P-labelled direct repeats (DR-1 to DR-5) as indicated in Fig. 9. The direct repeats were derived from the DR-5 element in the RAR-β2 promoter (de Thé et al., Nature, 343, 177-180, 1990) and modified to be separated by one to five nucleotides (Pettersson et al., Mechanisms of Dev., 54, 1-13, 1995). Protein-DNA complexes were separated on native 5% polyacryl-amide/0.25xTBE gels followed by autoradiography. As shown in Fig. 9, of the five DRs tested efficient VDRRγ binding could only be detected with DRs separated by three or four nucleotides and only in the presence of RXR. However, weaker RXR-dependent binding could also be observed to DR-2 and DR-1 elements. These

WO 99/19354 15 PCT/SE98/01548

results demonstrate that VDRRy require RXR heterodimerisation for efficient DNA-binding to a specific subset of DRs. These results, however, do not exclude the possibility that VDRRy may bind as a monomer, dimer or heterodimer to distinct but related DNA-sequences. Importantly, our results demonstrate that VDRRy and other nuclear receptors including the VDR (e.g. Markose, E. R. et al., Proc. Natl. Acad. Sci. USA, 87, 1701-1705, 1990), THRs (e.g. Gronemeyer, H. and Moras, D., Nature, 375, 190-191, 1995), LXRs (e.g. Willy, P. J. et al., Genes. Dev., 9, 1033-1045, 1995), have distinct but overlapping DNA-sequence and thus may regulate overlapping gene networks.

Interestingly, the most closely related nuclear receptor called ONR-1 (in Smith et al., 1994, Nucleic Acids Res., 22, pp66-71) or XOR-6 (in WO 96/22390) have been reported to "bind well to a retinoic acid response element, bRARE" (p. 11, line 30 in WO 96/22390). However, although the novel nuclear receptor VDRRg reported herein has 71% amino acid similarity in the DBD as compared to XOR-6 (fig 12), VDRRg does not appear to bind to the same bRARE sequence (DR-5 in Fig. 9).

15

20

25

30

10

5

EXAMPLE 6

Pregnenolone derivatives as activators of VDRRy

For identifying activators or ligands for VDRR γ , a library of substances structurally biased towards different classes of activators and ligands for nuclear receptors were tested. The activation of VDRR γ was analyzed in a reporter gene assay in transiently Caco-2 (TC7) cells (Carriere et al, 1994). In this initial screen, the synthetic substances with ability to activate VDRR γ were found to be structurally similar to pregnenolones (data not shown). Based on these results, naturally occuring pregnenolone derivatives were examined for activation of VDRR γ . The results are shown in Fig. 10. As is evident from Fig. 10, VDRR γ was activated about 5 to 12 fold by pregnenolone, 5 β -pregnane-3,20-dione, 5 β -pregnane-3 α ,20 β -diol and 3 α -hydroxy-5 β -pregnane-11,20-dione methanesulphonate. In contrast to the efficient activation observed by the 5 β -pregnane-3,20-dione, the corresponding planar steroid derivative 5 α -pregnane-3,20-dione did not activate the receptor. Other 5 β -pregnanes also activated VDRR γ efficiently as opposed to all planar pregnenolone derivatives tested, as is also evident from Fig. 10.

EXAMPLE 7

Pregnenolone 16 α -carbonitrile (PCN), dexamethasone and an antiprogestin (RU486) as activators of VDRR γ

Further experiments were performed to find out if pregnenolone 16α-carbonitrile (PCN), a glucocorticoid antagonist or dexamethasone are activators of VDRRγ. To this effect, Caco-2 cells were transfected as before with VDRRγ and the activation was analyzed after treatment of the cells with 10 μM PCN or dexamethasone. The results are shown in Fig. 11. As is evident from Fig. 11, VDRRγ was not activated by these substances, indicating that VDRRγ is not the human PCN receptor. This suggestion is corroborated by the observation that also the antiprogestin RU486 only caused a slight increase (two fold) in VDRRγ mediated reporter gene activity as is evident from Fig. 11.

Activators of XOR-6 (Fig. 3 in WO 96/22390) such as butyl 4-NH2 Benzoate did not activate VDRRγ (data not shown) in similar reporter assays as used in WO 96/22390.

SEQUENCE LISTING

<110> Pharmacia & Upjohn AB

<120> Novel vitamin D receptor related polypeptides, nucleic acid sequnce encoding the same and uses thereof.

<130> 1788 sequence listing

<140>

<141>

<150> 9703745-1

<151> 1997-10-14

<150> 9801148-9

<151> 1998-03-31

<160>4

<170> PatentIn Ver. 2.0

<210>1

<211>2910

<212> DNA

<213> Homo sapiens

<400> 1

cctctgaagg ttctagaatc gatagtgaat tcgtgggacg ggaagaggaa gcactgcctt 60 tacttcagtg ggaatctcgg cctcagcctg caagccaagt gttcacagtg aaaaaagcaa 120 gagaataage taataeteet gteetgaaca aggeagegge teettggtaa agetaeteet 180 tgatcgatcc tttgcaccgg attgttcaaa gtggacccca ggggagaagt cggagcaaag 240 aacttaccac caagcagtee aagaggeeea gaagcaaace tggaggtgag acceaaagaa 300 agetggaace atgetgaett tgtacaetgt gaggacacag agtetgttee tggaaagece 360 agtgtcaacg cagatgagga agtcggaggt ccccaaatct gccgtgtatg tggggacaag 420 gccactggct atcacttcaa tgtcatgaca tgtgaaggat gcaagggctt tttcaggagg 480 gccatgaaac gcaacgcccg gctgaggtgc cccttccgga agggcgcctg cgagatcacc 540 eggaagacce ggegacagtg ceaggeetge egeetgegea agtgeetgga gageggeatg 600 aagaaggaga tgatcatgtc cgacgaggcc gtggaggaga ggcgggcctt gatcaagcgg 660 aagaaaagtg aacggacagg gactcagcca ctgggagtgc agggggctgac agaggagcag 720 cggatgatga tcagggagct gatggacgct cagatgaaaa cctttgacac taccttctcc 780 catttcaaga atttccggct gccaggggtg cttagcagtg gctgcgagtt gccagagtct 840 ctgcaggccc catcgaggga agaagctgcc aagtggagcc aggtccggaa agatctgtgc 900 tetttgaagg tetetetgea getgegggg gaggatggea gtgtetggaa etacaaacce 960 ccagccgaca gtggcgggaa agagatette teeetgetge eccacatgge tgacatgtea 1020 acctacatgt tcaaaggcat catcagcttt gccaaagtca tctcctactt cagggacttg 1080

cccatcgagg accagatete cetgetgaag ggggccgett tegagetgtg teaactgaga 1140 ttcaacacag tgttcaacgc ggagactgga acctgggagt gtggccggct gtcctactgc 1200 ttggaagaca etgeaggtgg etteeageaa ettetaetgg ageceatget gaaatteeae 1260 tacatgetga agaagetgea getgeatgag gaggagtatg tgetgatgea ggecatetee 1320 ctettetece cagacegeee aggtgtgetg cageacegeg tggtggacea getgcaggag 1380 caattegeea ttaetetgaa gteetaeatt gaatgeaate ggeeceagee tgeteatagg 1440 ttettgttee gtgaagatea tggetatget eacegagete egeageatea atgeteagea 1500 cacccagegg etgetgegea tecaggacat acacccettt getaegeece teatgeagga 1560 gttgttcggc atcacaggta gctgagcggc tgcccttggg tgacacctcc gagaggcagc 1620 cagacccaga gecetetgag eegecaetee egggecaaga eagatggaca etgecaagag 1680 cegacaatge cetgetggee tgteteecta gggaatteet getatgacag etggetagea 1740 ttcctcagga aggacatggg tgcccccac ccccagttca gtctgtaggg agtgaagcca 1800 cagactetta egtggagagt geaetgaeet gtaggteagg accateagag aggeaaggtt 1860 gecettteet tttaaaagge eetgtggtet ggggagaaat eecteagate eeactaaagt 1920 gtcaaggtgt ggaagggacc aagcgaccaa ggataggcca tctggggtct atgcccacat 1980 acceaegttt gttegettee tgagtetttt eattgetace tetaatagte etgteteeca 2040 etteceacte gttecectee tetteegage tgetttgtgg geteaaggee tgtacteate 2100 ggcaggtgca tgagtatctg tgggagtcct ctagagagat gagaagccag gaggcctgca 2160 ccaaatgtca gaagettgge atgaceteat teeggeeaca teattetgtg tetetgeate 2220 catttgaaca cattattaag cactgataat aggtagcctg ctgtggggta tacagcattg 2280 actcagatat agatcctgag ctcacagagt ttatagttaa aaaaacaaac agaaacacaa 2340 acaatttgga tcaaaaggag aaaatgataa gtgacaaaag cagcacaagg aatttccctg 2400 tgtggatgct gagctgtgat ggcaggcact gggtacccaa gtgaaggttc ccgaggacat 2460 gagtetgtag gageaaggge acaaactgea getgtgagtg egtgtgtgtg atttggtgta 2520 ggtaggtctg tttgccactt gatggggcct gggtttgttc ctggggctgg aatgctgggt 2580 atgetetgtg acaaggetae getgacaate agttaaacae aceggagaag aaccatttae 2640 atgeaectta tattietgtg taeaeateta tteteaaage taaagggtat gaaagtgeet 2700 gccttgttta tagccacttg tgagtaaaaa tttttttgca ttttcacaaa ttatacttta 2760 tataaggcat tccacaccta agaactagtt ttgggaaatg tagccctggg tttaatgtca 2820 2910 aaaaaaaaa aaaaaaaaaa aaaaaagcnt

<210> 2

<211> 437

<212> PRT

<213> Homo sapiens

<400> 2

Met Glu Val Arg Pro Lys Glu Ser Trp Asn His Ala Asp Phe Val His
1 5 10 15

Cys Glu Asp Thr Glu Ser Val Pro Gly Lys Pro Ser Val Asn Ala Asp 20 25 30

Glu Glu Val Gly Gly Pro Gln Ile Cys Arg Val Cys Gly Asp Lys Ala 35 40 45

Thr Gly Tyr His Phe Asn Val Met Thr Cys Glu Gly Cys Lys Gly Phe 50 55 60

Phe Arg Arg Ala Met Lys Arg Asn Ala Arg Leu Arg Cys Pro Phe Arg 65 70 75 80

Lys Gly Ala Cys Glu Ile Thr Arg Lys Thr Arg Arg Gln Cys Gln Ala 85 90 95

Cys Arg Leu Arg Lys Cys Leu Glu Ser Gly Met Lys Lys Glu Met Ile 100 105 110

Met Ser Asp Glu Ala Val Glu Glu Arg Arg Ala Leu Ile Lys Arg Lys
115 120 125

Lys Ser Glu Arg Thr Gly Thr Gln Pro Leu Gly Val Gln Gly Leu Thr 130 135 140

Glu Glu Gln Arg Met Met Ile Arg Glu Leu Met Asp Ala Gln Met Lys
145 150 155 160

Thr Phe Asp Thr Thr Phe Ser His Phe Lys Asn Phe Arg Leu Pro Gly
165 170 175

Val Leu Ser Ser Gly Cys Glu Leu Pro Glu Ser Leu Gln Ala Pro Ser 180 185 190

Arg Glu Glu Ala Ala Lys Trp Ser Gln Val Arg Lys Asp Leu Cys Ser

195 200 205

Leu Lys Val Ser Leu Gln Leu Arg Gly Glu Asp Gly Ser Val Trp Asn 210 215 220

Tyr Lys Pro Pro Ala Asp Ser Gly Gly Lys Glu Ile Phe Ser Leu Leu 225 230 235 240

Pro His Met Ala Asp Met Ser Thr Tyr Met Phe Lys Gly Ile Ile Ser 245 250 255

Phe Ala Lys Val Ile Ser Tyr Phe Arg Asp Leu Pro Ile Glu Asp Gln 260 265 270

Ile Ser Leu Leu Lys Gly Ala Ala Phe Glu Leu Cys Gln Leu Arg Phe 275 280 285

Asn Thr Val Phe Asn Ala Glu Thr Gly Thr Trp Glu Cys Gly Arg Leu 290 295 300

Ser Tyr Cys Leu Glu Asp Thr Ala Gly Gly Phe Gln Gln Leu Leu Leu 305 310 315 320

Glu Pro Met Leu Lys Phe His Tyr Met Leu Lys Lys Leu Gln Leu His 325 330 335

Glu Glu Glu Tyr Val Leu Met Gln Ala Ile Ser Leu Phe Ser Pro Asp 340 345 350

Arg Pro Gly Val Leu Gln His Arg Val Val Asp Gln Leu Gln Glu Gln 355 360 365

Phe Ala Ile Thr Leu Lys Ser Tyr Ile Glu Cys Asn Arg Pro Gln Pro 370 375 380

Ala His Arg Phe Leu Phe Leu Lys Ile Met Ala Met Leu Thr Glu Leu 385 390 395 400

Arg Ser Ile Asn Ala Gln His Thr Gln Arg Leu Leu Arg Ile Gln Asp 405 410 415

Ile His Pro Phe Ala Thr Pro Leu Met Gln Glu Leu Phe Gly Ile Thr 420 425 430

Gly Ser Phe Ile Gly 435 <210>3

<211> 2802

<212> DNA

<213> Homo sapiens

<400> 3

tgaattcgtg ggcctgctgg gttagtgctg gcagcccccc tgaggccaag gacagcagca 60 tgacagteae caggacteae caetteaagg aggggteeet cagageaeet gecataceee 120 tgcacagtgc tgcggctgag ttggcttcaa accatccaag aggcccagaa gcaaacctgg 180 aggtgagacc caaagaaagc tggaaccatg ctgactttgt acactgtgag gacacagagt 240 ctgttcctgg aaagcccagt gtcaacgcag atgaggaagt cggaggtccc caaatctgcc 300 gtgtatgtgg ggacaaggcc actggctatc acttcaatgt catgacatgt gaaggatgca 360 agggettttt caggagggee atgaaaegea aegeeegget gaggtgeeee tteeggaagg 420 gegeetgega gateaecegg aagaeeegge gacagtgeea ggeetgeege etgegeaagt 480 gcctggagag cggcatgaag aaggagatga tcatgtccga cgaggccgtg gaggagaggc 540 gggccttgat caagcggaag aaaagtgaac ggacagggac tcagccactg ggagtgcagg 600 ggctgacaga ggagcagcgg atgatgatca gggagctgat ggacgctcag atgaaaacct 660 ttgacactac cttctcccat ttcaagaatt tccggctgcc aggggtgctt agcagtggct 720 gegagttgcc agagtetetg caggecccat egagggaaga agetgccaag tggagccagg 780 tccggaaaga tctgtgctct ttgaaggtct ctctgcagct gcggggggag gatggcagtg 840 tetggaacta caaaceecca geegacagtg gegggaaaga gatettetee etgetgeece 900 acatggetga catgteaace tacatgttea aaggeateat eagetttgee aaagteatet 960 cetaetteag ggaettgece ategaggaee agateteeet getgaagggg geegettteg 1020 agetgtgtca actgagattc aacacagtgt tcaacgcgga gactggaacc tgggagtgtg 1080 geoggetgte etactgettg gaagacaetg caggtggett ceageaactt etactggage 1140 ccatgctgaa attccactac atgctgaaga agctgcagct gcatgaggag gagtatgtgc 1200 tgatgcaggc catctccctc ttctccccag accgcccagg tgtgctgcag caccgcgtgg 1260 tggaccagct gcaggagcaa ttcgccatta ctctgaagtc ctacattgaa tgcaatcggc 1320 cccagcetge teataggite tigitectga agateatgge tatgeteace gageteegea 1380 gcatcaatge teageacace eageggetge tgegeateea ggacatacae ecetttgeta 1440

cgcccctcat gcaggagttg ttcggcatca caggtagctg agcggctgcc cttgggtgac 1500 accteegaga ggeageeaga eccagageee tetgageege eacteeeggg ecaagacaga 1560 tggacactgc caagageega caatgeeetg etggeetgte teeetaggga atteetgeta 1620 tgacagetgg ctageattee teaggaagga catgggtgee ecceaecee agtteagtet 1680 gtagggagtg aagccacaga ctcttacgtg gagagtgcac tgacctgtag gtcaggacca 1740 tcagagaggc aaggttgccc tttcctttta aaaggccctg tggtctgggg agaaatccct 1800 cagateceae taaagtgtea aggtgtggaa gggaceaage gaceaaggat aggeeatetg 1860 gggtctatgc ccacataccc acgtttgttc gcttcctgag tcttttcatt gctacctcta 1920 atagteetgt eteceaette ceaetegtte eceteetett eegagetget ttgtgggete 1980 aaggeetgta eteateggea ggtgeatgag tatetgtggg agteetetag agagatgaga 2040 agccaggagg cetgcaccaa atgtcagaag ettggcatga cetcatteeg gecacateat 2100 tctgtgtctc tgcatccatt tgaacacatt attaagcact gataataggt agcctgctgt 2160 ggggtataca gcattgactc agatatagat cctgagctca cagagtttat agttaaaaaa 2220 acaaacagaa acacaaacaa tttggatcaa aaggagaaaa tgataagtga caaaagcagc 2280 acaaggaatt teeetgtgtg gatgetgage tgtgatggea ggeaetgggt acceaagtga 2340 aggttcccga ggacatgagt ctgtaggagc aagggcacaa actgcagctg tgagtgcgtg 2400 tgtgtgattt ggtgtaggta ggtctgtttg ccacttgatg gggcctgggt ttgttcctgg 2460 ggctggaatg ctgggtatgc tctgtgacaa ggctacgctg acaatcagtt aaacacaccg 2520 gagaagaacc atttacatgc accttatatt tctgtgtaca catctattct caaagctaaa 2580 gggtatgaaa gtgcctgcct tgtttatagc cacttgtgag taaaaaatttt tttgcatttt 2640 cacaaattat actttatata aggcattcca cacctaagaa ctagttttgg gaaatgtagc 2700 cctgggttta atgtcaaatc aaggcaaaag gaattaaata atgtactttt ggctaaaaaa 2760 2802

<210>4

<211> 473

<212> PRT

<213> Homo sapiens

<400> 4

Met Thr Val Thr Arg Thr His His Phe Lys Glu Gly Ser Leu Arg Ala
1 5 10 15

- Pro Ala Ile Pro Leu His Ser Ala Ala Ala Glu Leu Ala Ser Asn His 20 25 30
- Pro Arg Gly Pro Glu Ala Asn Leu Glu Val Arg Pro Lys Glu Ser Trp 35 40 45
- Asn His Ala Asp Phe Val His Cys Glu Asp Thr Glu Ser Val Pro Gly 50 55 60
- Lys Pro Ser Val Asn Ala Asp Glu Glu Val Gly Gly Pro Gln Ile Cys 65 70 75 80
- Arg Val Cys Gly Asp Lys Ala Thr Gly Tyr His Phe Asn Val Met Thr 85 90 95
- Cys Glu Gly Cys Lys Gly Phe Phe Arg Arg Ala Met Lys Arg Asn Ala 100 105 110
- Arg Leu Arg Cys Pro Phe Arg Lys Gly Ala Cys Glu Ile Thr Arg Lys 115 120 125
- Thr Arg Arg Gln Cys Gln Ala Cys Arg Leu Arg Lys Cys Leu Glu Ser 130 135 140
- Gly Met Lys Lys Glu Met Ile Met Ser Asp Glu Ala Val Glu Glu Arg 145 150 155 160
- Arg Ala Leu Ile Lys Arg Lys Lys Ser Glu Arg Thr Gly Thr Gln Pro 165 170 175
- Leu Gly Val Gln Gly Leu Thr Glu Glu Gln Arg Met Met Ile Arg Glu 180 185 190
- Leu Met Asp Ala Gln Met Lys Thr Phe Asp Thr Thr Phe Ser His Phe 195 200 205
- Lys Asn Phe Arg Leu Pro Gly Val Leu Ser Ser Gly Cys Glu Leu Pro 210 215 220
- Glu Ser Leu Gln Ala Pro Ser Arg Glu Glu Ala Ala Lys Trp Ser Gln 225 230 235 240
- Val Arg Lys Asp Leu Cys Ser Leu Lys Val Ser Leu Gln Leu Arg Gly 245 250 255
- Glu Asp Gly Ser Val Trp Asn Tyr Lys Pro Pro Ala Asp Ser Gly Gly 260 265 270

Lys Glu Ile Phe Ser Leu Leu Pro His Met Ala Asp Met Ser Thr Tyr Met Phe Lys Gly Ile Ile Ser Phe Ala Lys Val Ile Ser Tyr Phe Arg Asp Leu Pro Ile Glu Asp Gln Ile Ser Leu Leu Lys Gly Ala Ala Phe Glu Leu Cys Gln Leu Arg Phe Asn Thr Val Phe Asn Ala Glu Thr Gly Thr Trp Glu Cys Gly Arg Leu Ser Tyr Cys Leu Glu Asp Thr Ala Gly Gly Phe Gln Gln Leu Leu Clu Pro Met Leu Lys Phe His Tyr Met Leu Lys Lys Leu Gln Leu His Glu Glu Glu Tyr Val Leu Met Gln Ala Ile Ser Leu Phe Ser Pro Asp Arg Pro Gly Val Leu Gln His Arg Val Val Asp Gln Leu Gln Glu Gln Phe Ala Ile Thr Leu Lys Ser Tyr Ile Glu Cys Asn Arg Pro Gln Pro Ala His Arg Phe Leu Phe Leu Lys Ile

Arg Leu Leu Arg Ile Gln Asp Ile His Pro Phe Ala Thr Pro Leu Met 450 455 460

Gln Glu Leu Phe Gly Ile Thr Gly Ser 465 470

CLAIMS

- A mammalian, preferably human, isolated or recombinant nucleic acid comprising a contiguous nucleic acid sequence encoding a vitamin D receptor related (VDRR) polypeptide.
- 2. An isolated or recombinant DNA/nucleic acid according to Fig. 1 or Fig. 7 or alleles thereof encoding a new VDRR polypeptide.
- 3. The nucleic acid according to claim 1 or claim 2 encoding the VDRR polypeptide containing a DNA-binding domain (DBD) comprising about 77 amino acids with 9 cysteine residues,, wherein said DBD is characterized by the following amino acid sequence similarity:
- (i) at least 60% amino acid sequence similarity with the DBD of hVDR; and
- (ii) at least 65% amino acid sequence similarity with the DBD of xONR1.
- 4. The nucleic acid according to claim 3, wherein said DBD is characterized by the following amino acid sequence similarity:
- (i) about 65% amino acid sequence similarity with the DBD of hVDR; and
- (ii) about 71% amino acid sequence similarity with the DBD of xONR1.
- 5. The nucleic acid according to any previous claim, encoding the VDRR polypeptide, wherein the ligand-binding domain (LBD) of said polypeptide is characterized by the following amino acid sequence similarity, relative to the LBDs of hVDR and xONR1, respectively:
- (i) at least about 30% amino acid sequence similarity with the LBD of hVDR; and
- (ii) at least about 40% amino acid sequence similarity with the LBD of xONR1.

- 6. The nucleic acid according to claim 5, wherein said LBD is characterized by the following amino acid sequence similarity:
- (i) at least 35% amino acid sequence similarity with the LBD of hVDR; and
- (ii) at least 45% amino acid sequence similarity with the LBD of xONR1.
- 7. The nucleic acid according to claim 6, wherein said LBD is characterized by the following amino acid sequence similarity:
- (i) about 42% amino acid sequence similarity with the LBD of hVDR; and
- (ii) about 54% amino acid sequence similarity with the LBD of xONR1.
- 8. The nucleic acid according to any previous claim, wherein said nucleic acid sequence is that given in Fig. 1 or Fig. 7 or alleles thereof.
- 9. The nucleic acid according to claim 8, wherein said nucleic acid sequence is the same or substantially the same as given in Fig. 1 or Fig. 7.
- 10. A nucleic acid probe for the detection of a nucleic acid sequence encoding a VDRR polypeptide in a sample.
- 11. The nucleic acid probe according to claim 10, wherein said probe comprises at least 14 contiguous nucleotides of the nucleic acid sequence given in Fig. 1 or Fig. 7.
- 12. A method for identifying clones encoding a VDRR polypeptide said method comprising screening a genomic or cDNA library with a nucleic acid probe according to claims 10 or 11 under low stringency hybridization conditions, and identifying those clones which display a substantial degree of hybridization to said probe.
- 13. An expression vector comprising a nucleic acid according to any of claims 1 -9.

- 14. A cell containing a nucleic acid according to any of claims 1 -9.
- 15. A cell containing an expression vector according to claim 14.
- 16. A process for recombinant production of a VDRR polypeptide, said process comprising expressing the nucleic acid of claims 1 to 9 in a suitable host cell.
- 17. The process according to claim 16, wherein the host cell is eukaryotic.
- 18. An isolated or recombinant mammalian, preferably human, VDRR polypeptide.
- 19. The isolated or recombinant VDRR polypeptide according to claim 18 comprising the amino acid sequence substantially the same or the same as given in Fig. 4 or Fig. 8.
- 20. A method to produce specific monoclonal and polyclonal antibodies to the polypeptide according to any of claims 18 and 19 comprising the injection of the protein to a mammalian.
- 21. A pharmaceutical formulation comprising an isolated or recombinant VDRR polypeptide according to claims 18 or 19, and one or more therapeutically acceptable excipients.
- 22. A method for identifying a ligand to a VDRR, by a cell-based reporter assay, transgenic-animal reporter assay or *in vitro*-binding assay.
- 23. A method for identifying a substance for treatment of a condition affected by a VDRR polypeptide, comprising screening for an agonist or an antagonist of VDRR polypeptide signal transduction to be used for treating metabolic, proliferative or inflammatory conditions.

- 24. A mammalian, preferably human, VDRR polypeptide according to any of claims 18 and 19 for use as a medicament.
- 25. Use of a substance affecting VDRR signal transduction, such as an agonist or an antagonist of VDRR polypeptide signal transduction, for the manufacture of a medicament for treating metabolic, proliferative or inflammatory conditions.
- 26. Use according to claim 25 of a substance affecting VDRR signal transduction for the manufacture of a medicament for treating obesity, diabetes, anorexia, lipoprotein defects, hyperlipidemia, hypercholesteremia or hyperlipoproteinemia.
- 27. Use of a substance affecting VDRR signal transduction for the manufacture of a medicament for treating osteoporosis, rheumatoid artritis, benign and malign tumors, hyperproliferative skin disorders or hyperparathyroidism.
- 28. Use according to any of claims 25-27, wherein the substance affecting VDRR signal transduction is a chemical molecule of natural or synthetic origin with a molecular weight in the range of from about 100 up to about 500 Da, preferably with a molecular weight of about 300 Da.
- 29. A method for treating metabolic, proliferative or inflammatory conditions comprising introducing into a mammal a nucleic acid vector according to claim 13 encoding for expression of a VDRR polypeptide and wherein said nucleic acid vector is capable of transforming a cell in vivo and expressing said polypeptide in said transformed cell.
- 30. A method for treatment of a metabolic, proliferative or inflammatory condition by administration of a therapeutically effective amount of a substance affecting VDRR signal transduction.

31. The method according to claim 30, wherein the substance affecting VDRR signal transduction is a chemical molecule of natural or synthetic origin with a molecular weight in the range of from about 100 up to about 500 Da, preferably with a molecular weight of about 300 Da.

International application No. PCT/SE 98/01548

		PCT/SE 9	18/01548		
A. CLASS	SIFICATION OF SUBJECT MATTER	·			
	CO7K 14/72, C12N 15/10, A61K 38/17 International Patent Classification (IPC) or to both nat S SEARCHED	tional classification and IPC			
	ocumentation searched (classification system followed by	classification symbols)			
	CO7K, C12N, A61K		· · · · · · · · · · · · · · · · · · ·		
	ion searched other than minimum documentation to the	extent that such documents are inclu	ded in the fields searched		
	FI,NO classes as above ata base consulted during the international search (name	of data have and where practicable	cearch terms used)		
Electronic da	ata oase consuled during the memational search (hame	of data base did, where processin,	······································		
C. DOCU	MENTS CONSIDERED TO BE RELEVANT				
Category*	Citation of document, with indication, where app	propriate, of the relevant passage	Relevant to claim No.		
X	WO 9622390 A1 (THE SALK INSTITUT STUDIES), 25 July 1996 (25.0 1	E FOR BIOLOGICAL 7.96), the claims, sec	1-19,21-22, 24-27		
A			20,23		
					
x	WO 9317041 A1 (THE GENERAL HOSPI 2 Sept 1993 (02.09.93), the	TAL CORPORATION), claims, seq. ID No 1	1-27		
					
X	WO 9636230 A1 (THE SALK INSTITUT STUDIES), 21 November 1996 (E FOR BIOLOGICAL 21.11.96)	1-27		
					
;					
X Furth	er documents are listed in the continuation of Box	C. X See patent family	annex.		
"A" docume	categories of cited documents: ent defining the general state of the art which is not considered	"T" later document published after date and not in conflict with th the principle or theory underly	the international filing date or priorit e application but cited to understand ing the invention		
"E" erlier d "L" docume	f particular relevance ocument but published on or after the international filing date ent which may throw doubts on priority claim(s) or which is	"X" document of particular relevance: the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "Y" document of particular relevance: the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art			
"O" docume means	establish the publication date of another citation or other reason (as specified) ent referring to an oral disclosure, use, exhibition or other				
	ent published prior to the international filing date but later than crity date claimed	"&" document member of the same	patent family		
Date of the	e actual completion of the international search	Date of mailing of the internati	_		
	ember 1998		12- 1998		
Swedish	mailing address of the ISA/ Patent Office	Authorized officer			
Box 5055	, S-102 42 STOCKHOLM No. + 46 8 666 02 86	Patrick Andersson Telephone No. +46 8 782 2	5 00		
	SA/210 (second sheet) (July 1992)				

International application No.
PCT/SE 98/01548

	PC1/30	90/01340
C (Continu	nation). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passa	ages Relevant to claim No.
X	WO 9306215 A1 (THE SALK INSTITUTE FOR BIOLOGICAL STUDIES), 1 April 1993 (01.04.93)	1-19,21-22
X	Nucleic Acids Research, Volume 22, No 1, 1994, Darrin P. Smith et al, "A novel nuclear receptor superfamily member in Xenopus that associates w RXR, and shares extensive sequence similarity to the mammalian vitamin D3 receptor" page 66 - page 71	ith
		
x	WO 8909223 A1 (ARCH DEVELOPMENT CORPORATION), 5 October 1989 (05.10.89), figure 5	1-19,21-22
P,X	EMBL, Databas Genbank/DDBJ, accession no. AF031814, Kliewer S.A. et al: "An orphan Nuclear Receptor Activated by Pregnanes Defines a Novel"; & Steroid Signaling Pathway, Cell 92:73-82(1998), 1-1709	1-27
	,	
	·	:
	1	
	,	

Form PCT/ISA/210 (continuation of second sheet) (July 1992)

International application No.

PCT/SE98/01548

Box I	Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)
This inte	ernational search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
i. 🛛	Claims Nos.: 28-30 because they relate to subject matter not required to be searched by this Authority, namely:
	See PCT Rule 39.1(iv): Methods for treatment of the human or animal
	body by surgery or therapy, as well as diagnostic methods.
_	
2	Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of Item 2 of first sheet)
This Inte	ernational Searching Authority found multiple inventions in this international application, as follows:
	g
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. 🔲	As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark	on Protest
	No protest accompanied the payment of additional search fees.

Information on patent family members

01/12/98

International application No. PCT/SE 98/01548

	tent document in search repor	1	Publication date		Patent family member(s)		Publication date
10	9622390	Al	25/07/96	AU	4693496	A	07/08/96
10	3022330	•••	40,00,00	CA	2210679		25/07/96
				EP	0822991	A	11/02/98
0	9317041	A1	02/09/93	AT	172247	T	15/10/98
	30270.2			AU	684458	В	18/12/97
				AU	3728393	A	13/09/93
				CA	2117491	A	02/09/93
				DE	69321580	D	00/00/00
				EP	0628054	A,B	14/12/94
				JP	7504416		18/05/95
				US	5686574	A	11/11/97
				US	5710017	A	20/01/98
				US	5756448	A	26/05/98
0	9636230	A1	21/11/96	UA	5426596	A	29/11/96
 10	9306215	A1	01/04/93	AU	2579692	A	27/04/93
•	3000220	•••	•=•	CA	2115452	A	01/04/93
				EP	0609240	A	10/08/94
				US	5571696	A	05/11/96
				US	5696233	A	09/12/97
				US	5710004	A	20/01/98
0	8909223	A1	05/10/89	EP	0407462	A	16/01/91
				JP	3504438	T	03/10/91
				US	5614620	_	25/03/97